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SITE SPECIFIC LISTERIA INTEGRATION VECTORS AND METHODS FOR USING THE SAME

ACKNOWLEDGMENT OF GOVERNMENT SUPPORT

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10 <u>CROSS-REFERENCE TO RELATED APPLICATIONS</u>

This application is a continuation-in-part of application serial no. 10/136,860 filed April 30, 2002; the disclosure of which is herein incorporated by reference.

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INTRODUCTION

Field of the Invention

The field of this invention is Listeria species, e.g., Listeria monocytogenes, particularly recombinant strains of Listeria species, and methods for their fabrication and use.

Background of the Invention

Listeria monocytogenes is a Gram-positive, food-borne human and animal pathogen responsible for serious infections in immunocompromised individuals and pregnant women. Severe L. monocytogenes infections in humans are characterized by meningitis, meningoencephalitis, septicemia, and fetal death. L. monocytogenes is ubiquitous in nature and, in addition, can be isolated from a wide variety of warm blooded animals.

Protocols for recombinantly engineering Listeria species are of interest in both research and therapeutic applications. For example, Listeria species transformation protocols find use in the elucidation of the mechanisms responsible for growth and virulence of these types of bacteria. In addition, such protocols also find use in the preparation of live Listeria vaccines, which vaccines find use in a variety of different medical applications.

To date, a variety of different protocols have been employed to transform Listeria species, where such protocols include: homologous recombination, transposon based recombination, etc. While different protocols are currently available for engineering Listeria species, such methods are not entirely satisfactory. Disadvantages currently experienced with one or more of the available protocols include: (1) instability of the expression cassette in the transformed species; (2) variable impact on virulence of the transformed species; (3) size constraints of the expression cassette that can be placed in the transformed species; etc.

As such, despite the variety of different transformation protocols available, there is continued interest in the identification of further transformation protocols that can expand the repertoire of available genetic tools. Of particular interest would be the development of an efficient, site-specific integration vector that was suitable for use with a wide array of different Listeria species, where the vector did not suffer from one or more of the above disadvantages of the currently available protocols.

Relevant Literature

Patents and published patent applications of interest include: U.S. Patent No. 5,830,702 and published PCT application serial nos.: WO 99/25376 and WO 00/09733.

SUMMARY OF THE INVENTION

Site-specific Listeria species integration vectors and methods for their use are provided. The subject vectors include a bacteriophage integrase gene and a bacteriophage attachment site, where in many embodiments the bacteriophage that is source of these elements is a listeriophage. In certain embodiments, the subject vectors further include a multiple cloning site, where the multiple cloning site may further include a coding sequence, e.g., a coding sequence for a heterologous polypeptide, etc. The subject vectors and methods find use in a variety of different applications, including the study of Listeria species and the preparation of Listeria species. vaccines.

BRIEF DESCRIPTION OF THE FIGURES

FIG. 1. (A) Plasmid map of the pPL1 integration vector. SEQ ID NO:24 provides the sequence of pPL1. Chloramphenicol resistance genes and *E. coli*

origin of replication are shown in grey, RP4 origin of transfer shown in white, integrase gene and *L. monocytogenes* p60 promoter shown in black. Multiple cloning site is shown at the bottom of the plasmid with unique restriction sites noted below the MCS in a box. pPL24 and pPL25 inserts are shown

5 schematically below the MCS and were cloned as described in Materials and Methods. Final sizes of the plasmid constructs and the restriction sites used in cloning are noted with each of the inserts. (B) Plasmid map of the integration vector pPL2. SEQ ID NO:28 provides the sequence of pPL2. The color scheme and genes are the same as in FIG. 1A except the PSA integrase and PSA*attPP* sites as noted. The multiple cloning site with 13 unique restriction sites is shown at the bottom of the plasmid.

- FIG. 2. Genomic organization of the attachment sites within the *comK* gene (A and B) and the tRNA^{Arg} gene (C and D). (A) Non-lysogenic *L. monocytogenes* strain, with an intact *comK* gene. Primers PL60 and PL61 amplify across the bacterial attachment site *comK-attBB'*. (B) Lysogenic *L. monocytogenes* strain (with approximately 40 kb of phage DNA inserted into the *comK* gene) or integrated strain (with pPL1 construct inserted into the *comK* gene). Primers PL14 and PL61 amplify across the hybrid attachment site *comK-attPB'*. (C) *L. monocytogenes* serotype 4b strain non-lysogenic at tRNA^{Arg}-attBB'.
- 20 Primers NC16 and NC17 amplify across the bacterial attachment site tRNA^{Arg}-attBB' in serotype 4b strains. Asterisk indicates primers NC16 and NC17 are substituted with PL102 and PL103 to amplify across the bacterial attachment site tRNA^{Arg}-attBB' in serotype 1/2 strains. (D) Lysogenic *L. monocytogenies* strain (with approximately 40 kb of phage DNA or 6 kb pPL2 vector DNA inserted at the 3' end of the tRNA^{Arg} gene. Primers NC16 and PL95 amplify across the hybrid attachment site tRNA^{Arg}-attBP' in both serotype 4b and 1/2 strains.
- FIG. 3. Expression and functional complementation of ActA in SLCC-5764. (A) Coomassie blue stained SDS-PAGE of SLCC-5764 derived strains grown to late-log phase. ActA is indicated by an arrow. Lane 1: molecular size marker; lane 2: DP-L3780; lane 3: DP-L4083; lane 4: DP-L4086; lane 5: SLCC-5764; lane 6: DP-L4082; lane 7: DP-L4084; lane 8: DP-L4085; lane 9: DP-L4087. Strains are described in Table 1. (B) Actin tail formation and movement of DP-

L4087 in *Xenopus* cell extract. The top panel is a phase image; the bottom panel is a fluorescent image of the same field.

FIG 4. (A) Clover-leaf diagram of tRNA^{Arg} utilized as the PSA attachment site. The arginine anticodon is circled. The region with sequence identity

5 between the tRNA gene and the PSA attPP' is outlined. The boundaries of the tRNA^{Arg} gene and Cove score (82.37) were predicted with tRNAscan-SE (31).

(B) Alignment of the tRNA^{Arg}-attBB' region of *L. monocytogenes* WSLC 1042 (top line) and the attPP' region of PSA downstream of the integrase gene. The 74 nt tRNA^{Arg} gene of *L. monocytogenes* is boxed and the 17 bp overlapping region of sequence identity (core integration site) is shaded grey. The tRNA^{Arg} gene anticodon is shown in bold and underlined. Identical nucleotide residues are indicated by (:). The numbers located at the left indicate the nucleotide position in the DNA sequences of the WSLC 1042 attachment site (AJ314913) and PSA genome (AJ312240).

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DESCRIPTION OF THE SPECIFIC EMBODIMENTS

Site-specific Listeria species integration vectors and methods for their use are provided. The subject vectors include a bacteriophage integrase gene and a bacteriophage attachment site, where in many embodiments the bacteriophage 20 that is source of these elements is a listeriophage. In certain embodiments, the subject vectors further include a multiple cloning site, where the multiple cloning site may further include a coding sequence, e.g., for a heterologous polypeptide, etc. The subject vectors and methods find use in a variety of different applications, including the study of Listeria species and the preparation of Listeria species vaccines.

Before the subject invention is described further, it is to be understood that the invention is not limited to the particular embodiments of the invention described below, as variations of the particular embodiments may be made and still fall within the scope of the appended claims. It is also to be understood that the terminology employed is for the purpose of describing particular embodiments, and is not intended to be limiting. Instead, the scope of the present invention will be established by the appended claims.

In this specification and the appended claims, the singular forms "a," "an" and "the" include plural reference unless the context clearly dictates otherwise. Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood to one of ordinary skill in the art to which this invention belongs.

Where a range of values is provided, it is understood that each intervening value, to the tenth of the unit of the lower limit unless the context clearly dictates otherwise, between the upper and lower limit of that range, and any other stated or intervening value in that stated range, is encompassed within the invention.

The upper and lower limits of these smaller ranges may independently be included in the smaller ranges, and are also encompassed within the invention, subject to any specifically excluded limit in the stated range. Where the stated range includes one or both of the limits, ranges excluding either or both of those included limits are also included in the invention.

Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood to one of ordinary skill in the art to which this invention belongs. Although any methods, devices and materials similar or equivalent to those described herein can be used in the practice or testing of the invention, representative methods, devices and materials are now described.

All publications mentioned herein are incorporated herein by reference for the purpose of describing various invention components that are described in the publications which might be used in connection with the presently described invention.

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In further describing the subject invention, the subject vectors are reviewed first in greater detail, followed by a review of the methods of using the subject vectors, as well as representative applications in which the subject vectors and methods find use.

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VECTORS

As summarized above, the subject invention provides Listeria site-specific integration vectors, i.e., vectors that integrate into Listeria genomes in a site-

specific manner. The subject vectors are characterized in that they stably integrate into a genome of a Listeria species in a site-specific, as opposed to random, manner. As the subject vectors integrate site-specifically into a target genome, the vectors of the subject invention typically are inserted into a specific 5 location or sequence (i.e., domain, location) of the target genome by a mechanims mediated by a recombinase, specifically an integrase, where the integrase is one that uses two distinct recognition sites as substrates, one of which is positioned in the integration site of the target genome (the site into which a nucleic acid is to be integrated) and another adjacent a nucleic acid of interest 10 to be introduced into the integration site (i.e., a site on the subject vector referred to herein as the "bacteriophage attachment site"). For example, the recognition sites for phage integrases are generically referred to as attB, which is present in the bacterial genome (into which nucleic acid is to be inserted) and attP (which is present in the phage nucleic acid adjacent the nucleic acid for insertion into the 15 bacterial genome, which is referred to herein as the "bacteriophage attachment site"). Recognition sites can be native (endogenous to a target genome) or altered relative to a native sequence. Use of the term "recognize" in the context of an integrase "recognizes" a recognition sequence, is meant to refer to the ability of the integrase to interact with the recognition site and facilitate site-20 specific recombination.

The subject vectors are capable of integrating into the genomes of a wide variety of different Listeria species. Integration is readily determined by using any convenient assay, including those known to those of skill in the art, that can identify whether or not vector DNA has integrated into the genomic DNA of a target organism. For example, vector DNA can be introduced into a target organism, e.g., via conjugation, and then integration can be determined via PCR amplification of the integrated sequence using primers flanking the target site, where the size of the amplification product is determinative of whether integration has occurred. An example of such an assay is provided in the experimental section, below. Additional features of many embodiments of the subject vectors is that they replicate autonomously in a non-Listeria host cell, e.g., E. coli, and are stable and innocuous in such non-Listeria host cells.

The subject integration vectors are typically double-stranded plasmid vectors, where the vectors are generally at least about 3 kb, often at least about 5

kb and may be large as 15 kb, 20 kb, 25 kb, 30 kb or larger, where the vectors sometimes range in size from about 3-6 to about 20 kb. The vectors include a number of structural features that impart to the vectors the above-described functional characteristics.

One structural feature of the subject vectors is a bacteriophage integrase gene, i.e., a nucleic acid coding sequence for a bacteriophage integrase, which is operably linked to a Listeria specific promoter, such that the gene (coding sequence) is expressed in the Listeria cell for which the vector is designed to be employed. In many embodiments, the bacteriophage integrase gene is one obtained from a listeriophage, i.e., it is a listeriophage integrase gene. A variety of different listeriophages are known in the art, where any convenient listeriophage may serve as the source of the integrase gene, i.e., the integrase encoding nucleic acid. Specific integrases of interest include, but are not limited to: the U153 integrase, the PSA integrase; and the like.

As indicated above, the integrase gene is operably linked to a promoter 15 (as well as regulatory and/or signal sequences, if necessary) that drives expression of the integrase gene when the vector is present in the Listeria cell for which the vector is designed and in which integration of the vector is desired. Any convenient promoter may be employed, where in certain embodiments the 20 promoter is a Listeria specific promoter. Representative promoters of interest include, but are not limited to: the Listeria p60 promoter, the Listeria actA promoter, the Listeria plcA promoter, the Listeria mpl promoter, the Listeria plcB promoter, the Listeria inIA promoter; a heat shock promoter; and the like Promoters may also include certain bacteriophage promoters such as the 25 promoters for T7, $Q\beta$, SP6 and the like if the strain of Listeria also expresses the cognate bacteriophage RNA polymerase. In addition to the integrase gene/promoter element described above, the subject vectors also include a phage attachment site, i.e., a sequence or domain of nucleotides that provide for a site-specific integration with a Listeria genome. Any convenient phage 30 attachment site may be employed, where selection of the phage attachment site will depend, at least in part, on the desired integration location for the vector. Representative phage attachment sites of interest include, but are not limited to: the comK integration site (as described in greater detail in the experimental

section below); the tRNA^{Arg} integration site (as described in greater detail in the experimental section below); and the like.

In addition, the subject vectors may include an origin of replication that provides for replication of the vector in a non-Listeria host cell, e.g., E.coli. This origin of replication may be any convenient origin of replication or ori site, where a number of ori sites are known in the art, where particular sites of interest include, but are not limited to: p15A; pSC101; ColEI; pUC; pMB9; and the like.

In addition, the subject vectors may include an origin of transfer site or element when convenient, e.g., when the vector is introduced in the target

10 Listeria cell using a conjugation protocol, as described in greater detail below. Any convenient origin of transfer (oriT) may be employed, where representative origins of transfer of interest include, but are not limited to: RP4 oriT; RSF1010 oriT;

and the like.

In addition, the subject vectors typically include at least one restriction endonuclease recognized site, i.e., restriction site, which is located on the vector at a location which is amenable to insertion of a heterologous gene/expression cassette. A variety of restriction sites are known in the art and may be present on the vector, where such sites include those recognized by the following restriction enzymes: HindIII, Pstl, Sall, Accl, HincII, Xbal, BamHI, Smal, Xmal, Kpnl, Sacl, EcoRI, BstXI, Eagl, Notl, Spel and the like. In many embodiments, the vector includes a polylinker or multiple cloning site, i.e., a closely arranged series or array of sites recognized by a plurality of different restriction enzymes, such as those listed above.

In certain embodiments, the subject vectors include at least one coding sequence, e.g., a coding sequence for heterologous polypeptide/protein coding sequence present in the multiple cloning site, e.g., as a result of using a restriction endonuclease site present in the multiple cloning site to insert the coding sequence into the vector, according to well known recombinant technology protocols. By "heterologous" is meant that the coding sequence encodes a product, e.g., a protein, peptide, polypeptide, glycoprotein, lipoprotein, or other macromolecule, that is not normally expressed in Listeria. In many embodiments, this coding sequence is part of an expression cassette, which provides for expression of the coding sequence in the Listeria cell for which the

vector is designed. The term "expression cassette" as used herein refers to an expression module or expression construct made up of a recombinant DNA molecule containing at least one desired coding sequence and appropriate nucleic acid sequences necessary for the expression of the operably linked coding sequence in a particular host organism, i.e., the Listeria cell for which the vector is designed, such as the promoter/regulatory/signal sequences identified above, where the expression cassette may include coding sequences for two or more different polypeptides, or multiple copies of the same coding sequence, as desired. The size of the coding sequence and/or expression cassette that includes the same may vary, but typically falls within the range of about 25-30 to about 6000 bp, usually from about 50 to about 2000 bp. As such, the size of the encoded product may vary greatly, and a broad spectrum of different products may be encoded by the expression cassettes present in the vectors of this embodiment.

The nature of the coding sequence and other elements of the expression 15 cassette may vary, depending on the particular application of the vector, e.g., to study Listeria species, to produce Listeria species vaccines, for cytosolic delivery of macromolecules, etc. For example, where the vectors are employed in the production of Listeria vaccines, the coding sequence may encode a heterologous. 20 antigen, where representative heterologous antigens of interest include, but are not limited to: (a) viral antigens, e.g., influenza np protein, HIV gag protein, HIV env protein or parts thereof, e.g., gp120 and gp41, HIV nef protein, HIV pol proteins, HIV reverse transcriptase, HIV protease, herpes virus proteins, etc., (b) malarial antigens; (c) fungal antigens; (d) bacterial antigens; (e) tumor and tumor 25 related antigens; and the like. Due to the flexibility of the vector system, virtually any coding sequence of interest may be inserted. Where secretion of the product encoded by the expression cassette is desired, the expression cassette may include a coding sequence for a fusion protein of a selected foreign antigen and a protein that directs secretion, e.g., Listeriolysin O or PI-PLC; a signal sequence, 30 such as hemolysin signal sequence, etc. Where the subject vectors are employed in the preparation of Listeria delivery vehicles, e.g., as described in PCT publication no. WO 00/09733 (the priority application of which is herein incorporated by reference); and Dietrich et al., Nature Biotechnology (1998) 16: 181-185, the heterologous polypeptide coding sequence may be a cytolysin, e.g.,

phospholipase, pore forming toxin, listeriolysin O, streptolysin O, perfringolysin O, acid activated cytolysins, phage lysins, etc. Other coding sequences of interest include, but are not limited to: cytokines, costimulatory molecules, and the like. As indicated above, the vector may include at least one coding sequence, where in certain embodiments the vectors include two or more coding sequences, where the coding sequences may encode products that act concurrently to provide a desired results.

In general, the coding sequence may encode any of a number of different products and may be of a variety of different sizes, where the above discussion merely provides representative coding sequences of interest.

METHODS OF PREPARING THE SUBJECT VECTORS

The vectors of the subject invention may be produced using any

convenient protocol, including by standard methods of restriction enzyme cleavage, ligation and molecular cloning. One protocol for constructing the subject vectors includes the following steps. First, purified nucleic acid fragments containing desired component nucleotide sequences as well as extraneous sequences are cleaved with restriction endonucleases from initial sources.

- 20 Fragments containing the desired nucleotide sequences are then separated from unwanted fragments of different size using conventional separation methods, e.g., by agarose gel electrophoresis. The desired fragments are excised from the gel and ligated together in the appropriate configuration so that a circular nucleic acid or plasmid containing the desired sequences, e.g. sequences corresponding
- 25 to the various elements of the subject vectors, as described above is produced. Where desired, the circular molecules so constructed are then amplified in a host, e.g. *E. coli*. The procedures of cleavage, plasmid construction, cell transformation and plasmid production involved in these steps are well known to one skilled in the art and the enzymes required for restriction and ligation are
- 30 available commercially. (See, for example, R. Wu, Ed., Methods in Enzymology, Vol. 68, Academic Press, N.Y. (1979); T. Maniatis, E. F. Fritsch and J. Sambrook, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y. (1982); Catalog 1982-83, New England Biolabs, Inc.; Catalog 1982-83, Bethesda Research Laboratories, Inc. An example of how to

construct a vector of the present invention is provided in the Experimental section, below.

METHODS

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Also provided by the subject invention are methods of using the above described Listeria site specific integration vectors to integrate a heterologous nucleic acid into a Listeria genome. In practicing the subject methods, a vector of the subject invention is introduced into a target Listeria cell under conditions sufficient for integration of the vector into the target cell genome to occur. Any convenient protocol for introducing the vector into the target cell may be employed.

Suitable protocols include: calcium phosphate mediated transfection; protoplast fusion, in which protoplasts harboring amplified amounts of vector are fused with the target cell; electroporation, in which a brief high voltage electric pulse is applied to the target cell to render the cell membrane of the target cell permeable to the vector; microinjection, in which the vector is injected directly into the cell, as described in Capechhi et al, Cell (1980) 22: 479; and the like. The above *in vitro* protocols are well known in the art and are reviewed in greater detail in Sambrook, Fritsch & Maniatis, Molecular Cloning: A Laboratory Manual (Cold Spring Harbor-Laboratory Press)(1989) pp16.30-16.55.

In certain embodiments, e.g., where direct introduction into the target
Listeria cell does not provide optimal results, one representative method that may
be employed is a conjugation method, which comprises: (a) introducing the

25 vector into a non-Listeria host cell, e.g., E.coli; followed by (b) transfer of the
vector from the non-Listeria host cell to the Listeria host cell, e.g., by conjugation.
Introduction into the non-Listeria host cell may be accomplished using any of the
protocols described above. For the conjugation step, any convenient protocol
may be employed, where suitable protocols typically include combining the donor

30 and acceptor cells in a suitable medium and maintaining under suitable
conditions for conjugation and vector transfer to occur. Specific representative
conditions are provided in the Experimental section below.

The above methods result in stable integration of the vector and any expression cassette carried thereby, e.g., that encodes a heterologous

protein/foreign antigen, into a target Listeria cell genome in a site specific manner. The subject methods find use with a wide variety of different Listeria species.

5 UTILITY

The above described vectors and methods of using the same find use in a variety of different applications, where representative applications include, but are not limited to: (a) research applications; (b) vaccine preparation applications; (c)

Listerial delivery vehicle preparation applications; and the like.

One type of application in which the subject vectors and methods of using may be employed is in research of Listeria species. For example, the subject vectors and methods allow simple and efficient strain construction and are widely useful in various strains used to study the intracellular life cycle of L.

15 monocytogenes. Additionally, the subject vectors and methods may be employed to produce stable merodiploid strains to allow refined copy number studies and studies of interactions within a protein through multimerization and testing of the dominance or recessive nature of different alleles of a gene in the same bacterial strain.

The subject vectors and methods also find use in vaccine preparation applications. For example, the subject vectors find use in the production of Listeria cultures capable of expressing a heterologous antigen, i.e., Listeria vaccines, where the Listeria cells employed may be attenuated. The attenuated strains employed may be capable of normal invasion of a host cell, but incapable of normal survival or growth in the cell or cell-to-cell spread, or they may have other alterations that preclude normal pathogenicity.

The vaccines produced using vectors of the present invention are administered to a vertebrate by contacting the vertebrate with a sublethal dose of the genetically engineered Listeria vaccine, where contact typically includes administering the vaccine to the host. Thus the present invention provides for vaccines genetically engineered with the integration vector and provided in a pharmaceutically acceptable formulation. Administration can be oral, parenteral, intranasal, intramuscular, intravascular, direct vaccination of lymph nodes, administration by catheter or any one or more of a variety of well-known

administration routes. In farm animals, for example, the vaccine may be administered orally by incorporation of the vaccine in feed or liquid (such as water). It may be supplied as a lyophilized powder, as a frozen formulation or as a component of a capsule, or any other convenient, pharmaceutically acceptable 5 formulation that preserves the antigenicity of the vaccine. Any one of a number of well known pharmaceutically acceptable diluents or excipients may be employed in the vaccines of the invention. Suitable diluents include, for example, sterile, distilled water, saline, phosphate buffered solution, and the like. The amount of the diluent may vary widely, as those skilled in the art will recognize. Suitable 10 excipients are also well known to those skilled in the art and may be selected, for example, from A. Wade and P.J. Weller, eds., Handbook of Pharmaceutical Excipients (1994) The Pharmaceutical Press: London. The dosage administered may be dependent upon the age, health and weight of the patient, the type of patient, and the existence of concurrent treatment, if any. The vaccines can be 15 employed in dosage forms such as capsules, liquid solutions, suspensions, or elixirs, for oral administration, or sterile liquid for formulations such as solutions or suspensions for parenteral, intranasal intramuscular, or intravascular use. In accordance with the invention, the vaccine may be employed, in combination with a pharmaceutically acceptable diluent, as a vaccine composition, useful in 20 immunizing a patient against infection from a selected organism or virus or with respect to a tumor, etc. Immunizing a patient means providing the patient with at least some degree of therapeutic or prophylactic immunity against selected pathogens, cancerous cells, etc.

The subject vaccines prepared with the subject vectors find use in

25 methods for eliciting or boosting a helper T cell or a cytotoxic T-cell response to a
selected agent, e.g., pathogenic organism, tumor, etc., in a vertebrate, where
such methods include administering an effective amount of the Listeria vaccine.
The subject vaccines prepared with the subject vectors find use in methods for
eliciting in a vertebrate an innate immune response that augments the antigen30 specific immune response. Furthermore, the vaccines of the present invention
may be used for treatment post-exposure or post diagnosis. In general, the use of
vaccines for post-exposure treatment would be recognized by one skilled in the
art, for example, in the treatment of rabies and tetanus. The same vaccine of the
present invention may be used, for example, both for immunization and to boost

immunity after exposure. Alternatively, a different vaccine of the present invention may be used for post-exposure treatment, for example, such as one that is specific for antigens expressed in later stages of exposure. As such, the subject vaccines prepared with the subject vectors find use as both prophylactic and therapeutic vaccines to induce immune responses that are specific for antigens that are relevant to various disease conditions.

The patient may be any human and non-human animal susceptible to infection with the selected organism. The subject vaccines will find particular use with vertebrates such as man, and with domestic animals. Domestic animals include domestic fowl, bovine, porcine, ovine, equine, caprine, Leporidate (such as rabbits), or other animal which may be held in captivity.

In general, the subject vectors and methods find use in the production of vaccines as described U.S. Patent No. 5,830,702, the disclosure of which is herein incorporated by reference; as well as PCT publication no W0 99/25376, the disclosures of the priority applications of which are herein incorporated by reference.

The subject vectors also find use in the production of listerial delivery vehicles for delivery of macromolecules to target cells, e.g., as described in: PCT publication no. WO 00/09733 (the priority application of which is herein incorporated by reference); and Dietrich et al., Nature Biotechnology (1998) 16: 181-185. A variety of different types of macromolecules may be delivered, including, but not limited to: nucleic acids, polypeptides/proteins, etc., as described in these publications.

KITS & SYSTEMS

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Also provided are kits and systems that find use in preparing the subject vectors and/or preparing recombinant Listeria cells using the subject vectors and methods, as described above. For example, kits and systems for producing the subject vectors may include one or more of: an initial vector with a multiple cloning site; a restriction endonuclease for cleaving a site in the multiple cloning site, a vector including an expression cassette of interest which is to be inserted into the multiple cloning site; etc. Where the kits and systems are designed for the production of the recombinant Listeria cells, the kits and systems may include

vectors, or components for making the same, as described above, Listeria target cells, non-Listeria host cells, and the like. The subject kits may further include other components that find use in the subject methods, e.g., reaction buffers, growth mediums, etc.

The various reagent components of the kits may be present in separate containers, or may all be precombined into a reagent mixture for combination with template DNA.

In addition to the above components, the subject kits will further include instructions for practicing the subject methods. These instructions may be present in the subject kits in a variety of forms, one or more of which may be present in the kit. One form in which these instructions may be present is as printed information on a suitable medium or substrate, e.g., a piece or pieces of paper on which the information is printed, in the packaging of the kit, in a package insert, etc. Yet another means would be a computer readable medium, e.g., diskette, CD, etc., on which the information has been recorded. Yet another means that may be present is a website address which may be used via the internet to access the information at a removed site. Any convenient means may be present in the kits.

The following examples are offered by way of illustration and not by way of limitation.

EXPERIMENTAL

25 I. MATERIALS AND METHODS

A. Construction of pPL1 integration vector.

Standard molecular techniques were used in the construction of the 6101 bp integration vector pPL1 (Fig. 1A). The complete sequence of the pPL1 vector is provided as SEQ ID NO:25. pPL1 is a low copy plasmid that replicates autonomously in *E. coli* and integrates in a site-specific manner in *L. monocytogenes*, and was assembled from 6 independent DNA sources as follows. Restriction sites in the PCR primers used for construction are

underlined. All PCR reactions used in cloning steps utilized Vent DNA polymerase (New England Biolabs).

The multiple cloning site (MCS) from pBluescript KS- (Alting-Mees, M. A., and J. M. Short. 1989. pBluescript II: gene mapping vectors. Nucleic Acids Res. 5 17:9494.) (bp 1-171) was cloned after PCR with primers 5'-

- GGACGTCATTAACCCTCACTAAAGG-3' and 5'-
- GGACGTCAATACGACTCACTATAGG-3' (SEQ ID NOS: 01 & 02). The low copy Gram-negative origin of replication and chloramphenicol acetyltransferase (CAT) gene from pACYC184 (Chang, A. C., and S. N. Cohen. 1978. Construction and
- 10 characterization of amplifiable multicopy DNA cloning vehicles derived from the P15A cryptic miniplasmid. J. Bacteriol. 134:1141-1156.) (bp 172-2253) were cloned after PCR with primers 5'-GGACGTCGCTATTTAACGACCCTGC-3' and 5'-GAGCTGCAGGAGAATTACAACTTATATCGTATGGGG-3' (SEQ ID NOS: 03 & 04). For direct conjugation from E. coli to L. monocytogenes, the RP4 origin of
- 15 transfer (oriT) (Pansegrau, W., E. Lanka, P. T. Barth, D. H. Figurski, D. G. Guiney, D. Haas, D. R. Helinski, H. Schwab, V. A. Stanisich, and C. M. Thomas. 1994. Complete nucleotide sequence of Birmingham IncP alpha plasmids. Compilation and comparative analysis. J. Mol. Biol. 239:623-663) (bp 2254-2624) was cloned from plasmid pCTC3 (Williams, D. R., D. I. Young, and M. Young.
- 20 1990. Conjugative plasmid transfer from Escherichia coli to Clostridium acetobutylicum. J. Gen. Microbiol. 136:819-826) after PCR with primers 5'-GCACTGCAGCCGCTTGCCCTCATCTGTTACGCC-3' and 5'-CATGCATGCCTCTCGCCTGTCCCCTCAGTTCAG-3' (SEQ ID NOS: 05 & 06). The listeriophage U153 integrase gene and attachment site (attPP') (A. Nolte, P.
- 25 Lauer, and R. Calendar, unpublished; bp 2629-4127)(SEQ ID NO: 25 includes the sequences for these sites), that direct the site-specific integration of the plasmid were cloned after PCR with primers 5'-GTAGATCTTAACTTTCCATGCGAGAGGAG-3' and 5'-

 - GGGCATGCGATAAAAAGCAATCTATAGAAAAACAGG-3' (SEQ ID NOS: 07 & -
- 30 08). For expression of the U153 integrase gene, the L. monocytogenes p60 promoter (Lauer, P., J. A. Theriot, J. Skoble, M. D. Welch, and D. A. Portnoy. 2001. Systematic mutational analysis of the amino-terminal domain of the Listeria monocytogenes ActA protein reveals novel functions in actin-based motility. Mol. Microbiol. 42:1163-1177.) was PCR amplified with primers 5'-

CCTAAGCTTTCGATCATCATAATTCTGTC-3' and 5'-

GGGCATGCAGATCTTTTTTTCAGAAAATCCCAGTACG-3' (SEQ ID NOS: 09 & 10) and cloned upstream of the integrase gene. Base pairs 4570-6101 are a *Hind* III-*Aat* II restriction fragment subcloned from pUC18-Cat (a kind gift from

5 Nancy Freitag), and contain the inducible Gram-positive CAT gene from pC194 (Horinouchi, S., and B. Weisblum. 1982. Nucleotide sequence and functional map of pC194, a plasmid that specifies inducible chloramphenicol resistance. J. Bacteriol. 150:815-825) (bp 4788-5850).

10 B. Cloning of the hly and actA genes into pPL1.

The *hly* gene was subcloned from the plasmid pDP-906 (Jones, S., and D. A. Portnoy. 1994. Characterization of *Listeria monocytogenes* pathogenesis in a strain expressing perfringolysin O in place of listeriolysin O. Infect. Immun. 62(12):5608-5613.) by restriction digestion with *Bam*H I and *Xba* I, gel purifying a

15 2.9 kb fragment and by ligating it into pPL1 cut with BamH I and Spe I. The resultant plasmid was designated pPL24 (Fig. 1A). The actA gene was PCR amplified from 10403S genomic DNA with primers 5'-

GGTCTAGATCAAGCACATACCTAG-3' and 5'-

CGGGATCCTGAAGCTTGGGAAGCAG-3'(SEQ ID NOS:11 & 12). The 2220 bp PCR product was gel purified, cut with *Bam*H I and *Xba* I, and cloned into pPL1 cut with *Bam*H I and *Spe* I. The resultant plasmid was designated pPL25 (Fig. 1A).

C. Phage curing, conjugation and molecular confirmation of plasmid integration.

Phage curing was accomplished by adapting historical methodologies (Cohen, D. 1959. A variant of Phage P2 Originating in *Escherichia coli*, strain B. Virology **7:**112-126; Six, E. 1960. Prophage substitution and curing in lysogenic cells superinfected with hetero-immune phage. J. Bacteriol. **80:**728-729.). *L.*

30 monocytogenes 10403S derivatives carrying a prophage at comK-attBB' (integrated in the comK ORF as described (Loessner, M. J., R. B. Inman, P. Lauer, and R. Calendar. 2000. Complete nucleotide sequence, molecular analysis and genome structure of bacteriophage A118 of Listeria monocytogenes: implications for phage evolution. Mol. Microbiol. 35(2):324-

340.)) were grown in BHI at 37°C to 108 CFU/ml and infected with listeriophage U153 (Hodgson, D. A. 2000. Generalized transduction of serotype 1/2 and serotype 4b strains of Listeria monocytogenes. Mol. Microbiol. 35(2):312-323.) at a multiplicity of infection of 20:1 in the presence of 5 mM CaCl₂. Cultures were 5 incubated by shaking at 37°C for 75 minutes and inhibition of growth was monitored by comparison of the OD_{600} of the infected culture with an uninfected control culture. The infected culture was diluted 10-2 and 10-4 in BHI, and both dilutions were grown at 37°C until the 10-2 dilution had increased 100-fold in optical density. The 10-4 fold dilution was then diluted 10-2, and 3 µl were plated 10 on BHI. Fifty colonies were tested for phage release initially by toothpicking colonies into 0.25 ml LB broth and replica plating at 30°C on a lawn of Mack-4R (DP-L862), a non-lysogenic rough strain of L. monocytogenes particularly susceptible to forming plaques. Candidates that did not plaque were then tested by spotting 10 µl of culture on a lawn of Mack-4R to detect plaque formation. If 15 this second test was negative, the candidate was tested whether it could support plaque formation by the phage from the parent 10403S strain (Φ 10403, (Hodgson, D. A. 2000. Generalized transduction of serotype 1/2 and serotype 4b strains of Listeria monocytogenes. Mol. Microbiol. 35(2):312-323.)). Curing was confirmed molecularly by PCR with the comK-attBB' specific primer pair 20 PL60/PL61 (sequences follow) for the absence of a phage at comK-attBB'. Approximately 10% of colonies were cured using this procedure.

Recipient strains of *L. monocytogenes* (SLCC-5764, DP-L1169 and DP-L1172) were made streptomycin resistant for counter-selection in conjugation experiments by plate selection on BHI supplemented with 200 µg/ml antibiotic.

pPL1 plasmid constructs were electroporated into *E. coli* strain SM10 (Simon, R., U. Priefer, and A. Pühler. 1983. A broad host range mobilization system for *in vitro* genetic engineering: transposon mutagenesis in Gram negative bacteria. Bio/Technology 1:784-791.) using standard techniques (Sambrook, J., T. Maniatis, and E. F. Fritsch. 1989. Molecular Cloning: A
 Laboratory Manual, 2nd edn. ed. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.). Bacterial strains were grown to mid-log (OD₆₀₀ ~0.55) shaking at 30°C. *E. coli* donor strains were grown in LB containing 25 μg/ml of chloramphenicol, *L. monocytogenes* recipient strains were grown in BHI. 2.5 ml

of donor culture was mixed with 1.5 ml of recipient culture and filtered onto prewashed 47 mm 0.45 µm HA type filers (Millipore). The filter was washed once with 10 ml BHI, transferred to a BHI plate with no antibiotics and incubated for 2 hours at 30°C. The bacterial cells were gently resuspended in 2.5 ml of BHI and 5 25 μl and 50 μl aliquots were plated in 3 ml of LB top agar on BHI plates supplemented with 7.5 μg/ml chloramphenicol and 200 μg/ml streptomycin. Plates were incubated at 30°C overnight and shifted to 37°C for 2-3 days. Individual colonies were picked and screened by PCR for integration at the phage attachment site using the primers PL14 (5'-CTCATGAACTAGAAAATGTGG-3') 10 (SEQ ID NO:13), PL60 (5'-TGAAGTAAACCCGCACACGATG-3')(SEQ ID NO:14) and PL61 (5'-TGTAACATGGAGGTTCTGGCAATC-3')(SEQ ID NO:15). PCR reactions were performed on small portions of individual bacterial colonies picked with sterile P200 pipet tips from BHI plates directly into 20 µI PCR reactions. The primer pair PL14/PL61 specifically amplifies attBP' in a PCR reaction, resulting in 15 a 743 bp product on integrated strains (both prophage and pPL1 derivatives). The primer pair PL60/PL61 specifically amplifies comK-attBB' in a PCR reaction, resulting in a 417 bp product only on non-lysogenic strains (i.e. DP-L4056). PCR assays were performed in a Hybaid Omn-E thermocycler with an annealing temperature of 55°C for 30 cycles. Integrants arose at a frequency of 20 approximately 2x10⁻⁴ per donor cell.

D. Hemolysis on blood plates and hemolytic activity assay.

Hemolysis on blood plates was scored on tryptic soy agar plates supplemented with 5% defimbrinated sheep blood (HemoStat, Davis CA).

25 Hemolytic assays were performed essentially as described (Portnoy, D. A., P. S. Jacks, and D. J. Hinrichs. 1988. Role of hemolysin for the intracellular growth of *Listeria monocytogenes*. J. Exp. Med. 167(4):1459-1471.). Hemolytic activity is expressed as the reciprocal of the dilution of culture supernatant required to lyse 50% of sheep erythrocytes.

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E. Plaquing in L2 cells.

Plaque sizes were determined as previously described (Lauer, P., J. A. Theriot, J. Skoble, M. D. Welch, and D. A. Portnoy. 2001. Systematic mutational

analysis of the amino-terminal domain of the *Listeria monocytogenes* ActA protein reveals novel functions in actin-based motility. Mol. Microbiol. 42(5):1163-1177.). Each strain was plaqued in 6 to 8 independent experiments and compared to 10403S in each experiment.

5

F. SDS-PAGE of surface expressed ActA.

Surface expressed ActA protein was prepared from late-log phase bacterial cultures grown in LB broth (OD₆₀₀ ~0.7) by resuspending equivalent amounts in SDS-PAGE buffer and boiling for 5 min. which extracts surface-expressed proteins but does not perturb the cell wall. Equivalent amounts were loaded on 7% SDS-PAGE and visualized with Coomassie blue.

G. Xenopus laevis cell extract motility assays.

X. laevis egg cytoplasmic extract was prepared as described (Theriot, J.

- 15 A., J. Rosenblatt, D. A. Portnoy, P. J. Goldschmidt-Clermont, and T. J. Mitchison. 1994. Involvement of profilin in the actin-based motility of *L. monocytogenes* in cells and in cell-free extracts. Cell 76(3):505-517.) and supplemented with tetramethylrhodamine iodoacetamide-labeled actin (Theriot, J. A., and D. C. Fung. 1998. *Listeria monocytogenes*-based assays for actin assembly factors.
- 20 Methods Enzymol. 298:114-122.). SLCC-5764-derived strains were grown overnight to stationary phase, washed 1×, added to cell extracts and incubated for 45 minutes before microscopic observation.

H. LD_{50} determinations.

Limited LD₅₀ were performed in BALB/c mice as described (Portnoy, D. A., P. S. Jacks, and D. J. Hinrichs. 1988. Role of hemolysin for the intracellular growth of *Listeria monocytogenes*. J. Exp. Med. 167(4):1459-1471.). Animal experiments were performed in the laboratory of Archie Bouwer at Oregon Health Sciences Center, Portland, OR.

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I. Identification of the PSA attachment site and construction of pPL2.

The PSA attachment site (tRNA^{Arg}-attBB') DNA sequence was obtained through a combination of inverse PCR and genome walking. Inverse PCR was

performed on Sau3 Al-digested DP-L4061 DNA (WSLC 1042 lysogenic for PSA) (SEQ ID NO:26 is the sequence of 2,072 bp surrounding WSLC 1042 tRNA^{Arg}-attBB') using the divergent primers PL95 (5'-

ACATAATCAGTCCAAAGTAGATGC)(SEQ ID NO:16) and PL97 (5'-

5 ACGAATGTAAATATTGAGCGG) (SEQ ID NO:17) that anneal within the PSA *int* gene. The resultant DNA sequence was used to design further oligonucleotides and these were used with the Genome Walker kit (Clontech), per the manufacturer's recommendations. DNA sequence and tRNA analysis was done with using MacVector (Accelrys), DNAsis (Hitachi), BLAST algorithm (2), and 10 tRNAscan-SE (Lowe, T. M., and S. R. Eddy. 1997. tRNAscan-SE: a program for improved detection of transfer RNA genes in genomic sequence. Nucleic Acids Res. 25(5):955-964.).

pPL1 was modified to utilize a different attachment site on the *L. monocytogenes* chromosome by replacing the U153 integrase gene and attachment site in the plasmid. The PSA *int* and *attPP'* were PCR amplified from PSA genomic DNA with primers PL100 (5'-GAAGATCTCCAAAAATAAACAGGTGGTGG) (SEQ ID NO:18) and PL101 (5'-CATGCATGCGTGGAGGAAAGAAGAAGAACGC) (SEQ ID NO:19)with Vent DNA polymerase, digested with *BgI* II and *Sph* I, and ligated to pPL1 that had been digested with the same enzymes. The resultant plasmid was designated pPL2 (FIG 1B). The sequence of pPL2 is provided as SEQ ID NO:28.

The DNA sequence of the PSA tRNA^{Arg}-attBB' from serotype 1/2 *L. monocytogenes* strains was obtained by a plasmid-trap strategy. DP-L4211 (pPL2 integrated in 10403S) genomic DNA was digested with *Nsi* I and *Nhe* I, which do not cleave in the vector, and ligated under dilute conditions to promote self-ligation. The ligations were transformed into XL1-blue and chloramphenicol resistant colonies were selected. The plasmids obtained were sequenced with the convergent primers PL94 (5'-GGAGGGAAAGAAGAACGC) (SEQ ID NO:20) and PL95 (sequence above) for attPB' and attBP', respectively, which flank attPP' in the PSA genomic DNA sequence. Further, because of the divergence between the sequences downstream of the tRNA^{Arg} gene among serotypes, a serotype 1/2 specific PCR assay across tRNA^{Arg}-attBB' was developed from the 10403S DNA sequence and used to determine the prophage status of various *L*.

monocytogenes strains. Primers PL102 (5'TATCAGACCTAACCCAAACCTTCC) (SEQ ID NO:21) and PL103 (5'AATCGCAAAATAAAAATCTTCTCG) (SEQ ID NO:22) specifically amplify a 533
bp PCR product in non-lysogenic serotype 1/2 strains. The primer pair NC16 (5'5 GTCAAAACATACGCTCTTATC)(SEQ ID NO:23) and PL95 specifically amplify a
499 bp PCR product in strains that are either lysogenic or contain an integration
vector at tRNA^{Arg}-attBB'. (SEQ ID NO:27 provides the 643 bp surrounding
10403S tRNA^{Arg}-attBB')

10 II. RESULTS AND DISCUSSION

A. pPL1 forms stable, single-copy integrants in various *L. monocytogenes* strains.

pPL1 is the first shuttle integration vector that we constructed to facilitate strain construction in *L. monocytogenes*. In order to test the pPL1 vector, we needed a *L. monocytogenes* strain that did not have a phage at the *comK* bacterial attachment site. We adapted historical methods to cure *L. monocytogenes* strains of their prophages and found after superinfection with phage U153, which has the same attachment site as the endogenous 10403S prophage, we were able to isolate prophage-free strains (see Materials and Methods). The prophage-cured 10403S strain was designated DP-L4056 and was used in subsequent experiments.

Conjugation was chosen as a method for introducing the vector into the target cells, as many Listeria spp. are inefficiently transformed. Conjugation of pPL1 from *E. coli* into *L. monocytogenes* was successful. Drug resistant transconjugants arose at a reproducible frequency of ~2x10⁻⁴ per donor *E. coli* cell, approximately three-fold lower than conjugation with autonomously replicating plasmids, indicating ~30% integration efficiency for strains receiving the plasmid by conjugation. All chloramphenicol resistant colonies were positive with the PCR assay using primers PL14 and PL61 and negative using a PCR assay across *attPP*' in pPL1 (PL14 paired with a primer in the RP4 *oriT*) indicating that they were true integrants and that the full genetic complement of plasmid pPL1 had integrated into the Listeria chromosome. In addition, this

experiment demonstrated that pPL1 was not maintained as an episomal plasmid and that the vector did not integrate as a concatamer.

We tested the stability of the integrants under non-selective growth conditions. Three integrant strains, DP-L4074 and the merodiploid strains DP-L4076 and DP-L4078 (described in the following sections) were passed in liquid BHI media for 100 generations and plated for single colonies. Ninety-six colonies were then exposed to 0.1 µg/ml chloramphenicol (to induce CAT gene expression) and patched on plates containing 7.5 µg/ml antibiotic. All colonies retained drug resistance. Thirty colonies from each non-selective growth experiment were assayed with the PL14/PL61 PCR assay and all PCR reactions resulted in the 743 bp product, indicating all transconjugants retained the integrated plasmid.

We further addressed whether the integration vector would be generally useful for any *L. monocytogenes* strain with an available attachment site. There have been greater than 320 listeriophages isolated, and many have restricted host ranges. It was unclear whether there was a biological barrier to U153 integrase gene function in host strains that do not support U153 infection. We therefore picked three strains that did not contain a prophage at the *comK* attachment site; two serotype 4b clinical isolates and SLCC-5764, a serotype 1/2a strain that constitutively expresses the known virulence factors in an unregulated manner and has been useful for studying these virulence factors *in vitro*. Each of these strains was first made streptomycin resistant for counter selection in conjugation experiments (as described in Materials and Methods). Streptomycin resistant derivatives were chosen on the basis of having the same 25 growth rate as the parent strain to avoid experimental complications related to viability. The resultant strains, DP-L4088, DP-L4089, and DP-L4082 all proved suitable recipients for pPL1 integration at a similar frequency to DP-L4056.

Finally, we conducted a survey of *L. monocytogenes* isolates to identify suitable strains that do not harbor a prophage at the *comK* attachment site using the PCR assays across *comK-attBB*' (primers PL60/PL61) and the hybrid *attPB*' (primers PL14/PL61). The results of these experiments (Table 2) indicated many of the strains commonly used to study the biology and pathogenesis of *L. monocytogenes* including 10403S, L028, and EGDe had a prophage at *comK*.

Table 2. Prophage status of various strains

Strain	Description	Source	serotype	PL60/PL61	PL14/PL61
10403S	wild type			comK	attPB'
DP-L4056	_	rabbit pellets	1/2a	_ a	+
	10403S phage cured	This work	1/2a	+ b	-
DP-L861	SLCC-5764 (Mack)	WT (overexpresser)	1/2a	+	_
DP-L3818	WSLC 1118::A118	Camembert cheese	4b	<u>-</u>	+
DP-L3633	EGDe	WT (1960s human)	1/2a	-	+
DP-L3293	LO28	WT (clinical origin)	1/2c	_	+
DP-L185	F2397	L.A., Jalisco cheese	4b	+	•
DP-L186	ScottA	Massachusettes	4b	+	-
		outbreak, milk		•	-
DP-L188	ATTC 19113	Denmark, human	3	+	
DP-L1168	clinical	cole slaw	4b	, +	-
DP-L1169	clinical	patient	4b	+	-
DP-L1170	clinical	patient	4b	+	-
DP-L1171	clinical	brie	1/2b	+	-
DP-L1172	clinical	alfalfa tablets	4b	+	-
DP-L1173	clinical	deceased patient	4b	T	-
DP-L1174	clinical	deceased patient	4b	-	+
DP-L3809	1981 Halifax	placenta	4b	-	+
DP-L3810	1981 Halifax	CSF & brain	4b	+	-
DP-L3812	1981 Halifax	coleslaw	4b	+	-
DP-L3813	1996 Halifax	blood	40 ?	+	-
DP-L3814	1981 Halifax	CSF	<i>:</i> 4b	-	+
DP-L3815	1993 Halifax	CSF	46 1/2a	+	-
DP-L3816	1995 Halifax	blood	1/2a ?	+	-
OP-L3817	1993 Halifax	CSF		+	-
OP-L3862	1998 Michigan	patient	1/2a	+	-
	o DCD manula C	pair noted at top of colu	4b	- POD	+

a (-): negative PCR result for primer pair noted at top of column.
 b (+): positive PCR result for primer pair noted at top of column. The PL60/PL61 primer pair specifically amplify a 417 bp PCR product in non-lysogenic strains and result in no PCR product in lysogenic strains. The PL14/PL61 primer pair specifically amplify a 743 bp PCR product in lysogenic strains and result in no PCR product in non-lysogenic strains.

B. The status of comK did not affect the virulence of L. monocytogenes.

We next compared DP-L4056 and DP-L4074 to wild-type 10403S in standard virulence assays to determine if the presence of a prophage at *comK*, lack of prophage, or integration vector altered the virulence phenotypes. These three strains were assayed for LLO activity, ability to form plaques in monolayers of L2 cells, and for virulence in the mouse LD₅₀ assay (Table 3). All were indistinguishable from one another, strongly suggesting that the integrity of the *comK* ORF and the presence of pPL1 had no measurable impact on virulence.

Table 3. Complementation of actA and hly

Strain	Genotype	Hemolysis on blood plates	Hemolytic activity ^a	Plaque size ^b	LD ₅₀ ^c
10403S	wild type	+	nd	100.(na)	$\sim 2 \times 10^4$
DP-L4056	10403S phage cured	+	97	101 (1.4)	$<1 \times 10^{5}$
DP-L4074	DP-L4056 comK::pPL1	+	98	99 (1.4)	$<1x10^5$
DP-L4027	DP-L2161 phage cured, Δhly	-	0	0 (0)	1x10 ⁸
DP-L4075	DP-L4027 Δhly, comK::pPL24	+	99	97 (3.9)	<1x10
DP-L4076	DP-L4056 comK::pPL24	+	198	96 (2)	nd
DP-L4029	DP-L3078 phage cured, ΔactA	nd	nd	0 (0)	2x10 ⁷
DP-L4077	DP-L4029 ΔactA, comK::pPL25	nd	nd	86 (4)	<1x10
DP-L4078	DP-L4056 comK::pPL25	nd	nd	72 (6.8)	<1X10 nd

 ^a Hemolytic units data shown is from one representative experiment. nd: not determined. ^b Plaque size is the average of 8 to 10 independent experiments and shown as a percent of wild type (defined as 100%). Standard deviations are shown in parentheses. na: not applicable. ^c LD₅₀s of 10403S and Δ*hly* (DP-L2161) were determined in (37), the LD₅₀ of the Δ*actA* strain (DP-L1942, a smaller deletion within the *actA* ORF that does not support actin nucleation at the bacterial surface) was determined in (4).

10

C. Full complementation of hly at the phage attachment site.

Listeriolysin-O (LLO), the gene product of *hly*, is a secreted pore-forming cytolysin that is responsible for escape from the membrane-bound vacuole when *L. monocytogenes* first enters a host cell. LLO is absolutely required for the intracellular life cycle of *L. monocytogenes* and virulence. LLO activity can be measured by hemolytic activity on red blood cells. *Hly* mutants fail to form plaques in monolayers of L2 cells and are 5 logs less virulent in the mouse LD₅₀ assay.

We cloned the *hly* structural gene into pPL1 and conjugated this plasmid from *E. coli* to phage-cured wild type and Δ*hly L. monocytogenes* derivatives, resulting in DP-L4076 (an *hly* merodiploid) and DP-L4075 (*hly* only at the phage *comK-att* site). These strains were tested for hemolytic activity on blood plates, for the relative amount of hemolytic units secreted, ability to form a plaque in a

monolayer of L2 cells, and virulence in the mouse LD₅₀ assay (Table 3). The quantitative complementation of *hly* in the deletion strain background and the doubling of hemolytic units produced in the merodiploid strain indicate two things. First, gene expression is not *de facto* affected by ectopic expression at the *comK* chromosomal position. Second, the *hly* promoter is self-contained. Additionally, a two-fold increase in the amount of LLO is not deleterious to the virulence and intracellular life cycle of *L. monocytogenes*, at least as measured by plaquing.

D. Complementation of actA at the phage attachment site approaches wild-type expression.

10 ActA, a second major *L. monocytogenes* virulence factor, is responsible for commandeering host cell actin-cytoskeletal factors used for intracellular bacterial motility. ActA is also absolutely required for bacterial pathogenesis as mutants in *actA* are both unable to spread from cell-to-cell and form a plaque in a cell monolayer and are 3 logs less virulent than wild type. Additionally, ActA expression appears to be more complex than that of LLO: there are two promoters that drive *actA* expression. One is immediately upstream of the *actA* ORF and the second is in front of the *mpl* gene upstream of *actA*.

We constructed several strains to evaluate the complementation of *actA* at the phage attachment site. The first group included making second-site

20 complemented (DP-L4077) and merodiploid (DP-L4078) strains in the 10403S background. These were assayed for plaque formation in an L2 monolayer (Table 3). Integrated ActA did not fully complement in this assay (plaque size of 86%) and the merodiploid strain formed an even smaller plaque (72% of wild type). We interpret these results to indicate that there may be a small contribution of the second promoter upstream of the *mpl* gene for optimal *actA* expression. Additionally, there appears to be a critical concentration of ActA on the surface of intracellular bacteria because two copies of *actA* (with 3 promoters driving expression) further decreases the ability to spread from cell to cell, presumably because there is too much ActA on the bacterial surface for optimal motility.

We further tested ActA complementation in the virulence gene over-expressing strain SLCC-5764. ActA is effectively expressed in this strain from the *comK-attBB*' site (Fig 3A, lane 9). Considering the plaquing data of 10403S

complemented *actA* strains, it may have been predicted that the merodiploid strain DP-L4085 would make more ActA than the parent strain. However, this was not observed: the parent strain, the complemented strain and the merodiploid strain all expressed similar levels of ActA (Fig 3A, lanes 5, 8, and 9).

- 5 This observation was likely due to the complete lack of regulation and high level of constitutive expression of ActA in SLCC-5764. Additionally, DP-L4087 supports actin nucleation at the bacterial surface, actin tail formation and bacterial motility in cell extracts (Fig. 3B). The results of these cell-extract experiments indicate that the integration vector system for complementation will
- be useful for *in vitro* studies of *L. monocytogenes* motility, facilitating strain construction and placing various molecular constructs in different host strains for study in a desired set of assays. In particular, several alleles of *actA* that have unusual motility phenotypes have been transferred to the SLCC-5764 Δ*actA* strain using pPL1 and are currently being evaluated in cell extracts. The study of
- 15 these mutants in the simplified cell-extract system should yield insights into the activities of poorly understood regions of the ActA protein.

The strains referenced above are provided in the following Table 1.

Strain	Relevant genotype or plasmid
E. coli:	
SM10	Conjugation donor. F- thi-1 thr-1 leuB6 recA tonA21 lacYl
	$supE44 Mu+C l- (RP4-2(Tc::Mu)1 Km^r Tra+$
XL1-Blue	Plasmid manipulations. recAl endAl gyrA96 thi-1 hsdR17
	SupE44 relA1 lac [F'proAB lac1 Z \DM15 Tn10 (Tet)]
DP-E4067	integration vector pPL1/SM10
DP-E4068	hly integration vector pPL24/SM10
DP-E4069	actA integration vector pPL25/SM10
DP-E4190	Integration vector pPL2/SM10
<u>L. monocytogene</u>	<u>es:</u>
10403S	wild type
DP-L4056	10403S phage cured
OP-L4027	DP-L2161 phage cured, Δhly
DP-L4029	DP-L3078 phage cured, ΔactA
DP-L4074	DP-L4056 comK::pPL1
OP-L4075	DP-L4027 Δhly, comK::pPL24
DP-L4076	DP-L4056 comK::pPL24
DP-L4077	DP-L4029 ΔactA, comK::pPL25
OP-L4078	DP-L4056 comK::pPL25
LCC-5764	Virulence gene over expresser (Mack, DP-L861)
P-L862	Mack-4R (SLCC-5764 rough isolate)
P-L4082	SLCC-5764 Str' derivative
P-L3780	SLCC-5764 $\triangle actA$ (deletion of amino acids 7-633)

Strain	Relevant genotype or plasmid	
DP-L4083	DP-L3780S (Str' dervivative)	
DP-L4084	DP-L4082, comK::pPL1	
DP-L4085	DP-L4082, comK::pPL25	
DP-L4086	DP-L4083, comK::pPL1	
DP-L4087	DP-L4083 ΔactA, comK::actA	
DP-L4088	DP-L1169S 4b strain, Str'	
DP-L4089	DP-L1172S 4b strain, Str	
DP-L4090	DP-L4088, comK::pPL1	
DP-L4091	DP-L4089, comK::pPL1	
DP-L4199	EGDe, Str' derivative	
DP-L4026	WSLC 1042, (ATCC 23074)	
DP-L4061	WSLC 1042::PSA	
DP-L4221	10403S, tRNA Arg::pPL2	

E. Phage PSA integrates into a tRNA^{Arg} gene and pPL2 construction.

pPL1 integration into L. monocytogenes strains that harbour a prophage in the comK attachment site is hindered by the process of first having to cure the 5 prophage from the host strain. To alleviate the need for the phage-curing step, the specificity of pPL1 integration was changed to that of the PSA prophage. PSA, (Phage from ScottA) is the prophage of L. monocytogenes strain ScottA, a serotype 4b strain that was isolated during an epidemic of human listeriosis. Using the PSA genomic DNA sequence, we identified an integrase-like ORF with 10 a contiguous non-coding sequence that we predicted to contain the attPP' sequences. The PSA integrase sequence was then used to obtain the DNA sequence of PSA-attBB' from the PSA lysogenic strain DP-L4061 (see Materials and Methods). PSA was found to integrate in a tRNA^{Arg} gene that is 88% identical to a tRNA gene (trnSL-ARG2) from B. subtilis. The anticodon of the 15 tRNA^{Arg} gene is 5'UCU, the most commonly used arginine anticodon in *L*. monocytogenes. The PSA and bacterial attachment sites share 17 bp of DNA identity, and the tRNA Arg-attBB' contains a short nucleotide sequence that completes the tRNA^{Arg} sequence that is interrupted by integration of PSA (Fig 4). The attachment site $tRNA^{Arg}$ gene is present only once in the genome of L. 20 monocytogenes strain EGDe and apparently only once in the serotype 4b. This indicates that not only is the PSA integration site unique, but also that precise reconstitution of the gene upon integration (or excision) is likely required for survival of the cell.

pPL2 was constructed by replacing the U153 listeriophage integrase gene and attachment site in pPL1 with the PSA listeriophage integrase gene and attachment site. pPL2 was tranformed into SM10 and the resulting strain was mated into 10403S, EGDe (carrying a streptomycin resistance mutation) and the 5 serotype 4b strain DP-L4088. Chloramphenicol resistant transconjugants arose from each of these crosses at approximately 2x10⁻⁴ per donor cell, the same rate as pPL1 integration. Two recombinants from each cross were restreaked under drug selection and tested by PCR for the presence of PSA-attBP' using primers NC16 and PL95. The expected 499 bp PCR product was obtained in each of the 10 colonies tested, indicating pPL2 integrates into tRNA^{Arg}-attBB' in both serotype 1/2 and 4b strains. We tested the stability of the integrated pPL2 in both EGDe and DP-L4088 strains with the same non-selective 100-generation experiment described for pPL1. Forty-nine colonies from each of the amplified cultures were tested for chloramphenicol resistance. The EGDe-derived strains retained 100% 15 drug resistant colonies indicating complete stability of the integrants. In the case of the DP-L4088 integrant, two of the 49 colonies were chloramphenicol sensitive, suggesting a low level of excision can occur in this serotype 4b strain. In order to test whether precise excision had occured, we PCR amplified across tRNA^{Arg}-attBB' and sequenced the PCR products. The wild-type DNA sequence 20 was obtained, indicating a precise excision event.

During the course of our PCR experiments, we noted a divergence between the tRNA^{Arg}-attPB' sites from serotype 4b and serotype 1/2 L. monocytogenes. To determine nature of this divergence, we isolated and sequenced the tRNA^{Arg}-attBB' site from 10403S (as described in Materials and 25 Methods). We found that the sequence of attPB' in 10403S (3' of the tRNA^{Arg} gene) is unrelated to that of the serotype 4b strain WSLC 1042. In contrast to this, the sequence of attBP' (5' of the tRNA^{Arg} gene) in 10403S is 96-97% identical to the corresponding regions in L. monocytogenes serotype 4b strain WSLC 1042, the serotype 4b strain sequenced by TIGR, and the serotype 1/2a strain EGDe sequenced by the European Listeria Consortium. Thus, the bacterial attBB' sequences recognized by the PSA integrase are likely to encompass more of the attB DNA sequence than the attB' DNA sequence. Additionally, we tested the availability of the tRNA^{Arg}-attBB' in the common

laboratory strains of *L. monocytogenes* with a PCR assay using primers PL102 and PL103. We found the tRNA^{Arg} attachment site to be available in strains 10403S, EGEe, and L028 indicating that pPL2 may be readily utilized in these backgrounds for strain construction, complementation, and genetics studies without first curing endogenous prophages.

F. <u>Expression of Aquoria victoria GFP</u>.

A GFP coding sequence as described in United States Patent No. 5,777,079 (the disclosure of which is herein incorporated by reference), is cloned 10 into plasmid pPL1, transferred into the genome of L. monocytogenes. The GFP coding sequence is amplified by polymerase chain reaction (PCR) and the PCR fragment is cloned into the multiple cloning site of pPL1. A suitable promoter, containing appropriate transcriptional elements and a translational leader sequence for expressing the GFP in L. monocytogenes are cloned at the 5' end 15 of the GFP coding sequence such that they induce the expression of the GFP protein. The modified pPL1 plasmid constructs are electroporated into E. coli strain SM10 using standard techniques, and the modified pPL1 plasmid construct is conjugated into L. monocytogenes as described above. Recombinant L. monocytogenes are selected on BHI plates supplemented with 7.5 µg/ml 20 chloramphenicol and 200 μg/ml streptomycin. Individual colonies are picked and screened by PCR for integration at the phage attachment site using the primers PL14 (5'-CTCATGAACTAGAAAAATGTGG-3') (SEQ ID NO:13), PL60 (5'-TGAAGTAAACCCGCACACGATG-3')(SEQ ID NO:14) and PL61 (5'-TGTAACATGGAGGTTCTGGCAATC-3')(SEQ ID NO:15). Cultures of 25 recombinant *L. monocytogenes* are grown, prepared and screened for GFP.

III. Utility of pPL1 and pPL2.

The construction and characterization of the first single step site specific integration vectors for use in *L. monocytogenes* furthers the genetic tools available for the study of this pathogen. These vectors allow more facile strain construction than historic methods and are widely useful in various strains used to study the intracellular life cycle of *L. monocytogenes*. Additionally, stable merodiploid strains can be constructed to allow refined copy number studies and

studies of interactions within a protein through multimerization and testing of the dominance or recessive nature of different alleles of a gene in the same bacterial strain.

pPL1 and pPL2 are also useful for vaccine development, e.g., for at least 5 enhancing, including both eliciting and boosting, an immune response to a target cells or cells, e.g., a foreign pathogen, etc. Several recombinant L. monocytogenes systems have been used to elicit cell-mediated immune responses in mice (Frankel, F. R., S. Hegde, J. Lieberman, and Y. Paterson. 1995. Induction of cell-mediated immune responses to human immunodeficiency 10 virus type 1 Gag protein by using Listeria monocytogenes as a live vaccine vector. J. Immunol. 155(10):4775-4782.; Goossens, P. L., G. Milon, P. Cossart, and M. F. Saron. 1995. Attenuated Listeria monocytogenes as a live vector for induction of CD8+ T cells in vivo: a study with the nucleoprotein of the lymphocytic choriomeningitis virus. Int. Immunol. 7(5):797-805; Ikonomidis, G., Y. 15 Paterson, F. J. Kos, and D. A. Portnoy. 1994. Delivery of a viral antigen to the class I processing and presentation pathway by Listeria monocytogenes. J. Exp. Med. 180(6):2209-2218.; Shen, H., M. K. Slifka, M. Matloubian, E. R. Jensen, R. Ahmed, and J. F. Miller. 1995. Recombinant Listeria monocytogenes as a live vaccine vehicle for the induction of protective anti-viral cell-mediated immunity. 20 Proc. Natl. Acad. Sci. USA 92(9):3987-3991.). One limitation with plasmid-based expression of recombinant proteins in L. monocytogenes is the stability of the plasmids in vivo (i.e. in the host animal) without selection. Additionally, chromosomal construction of strains expressing foreign antigens is time consuming. pPL1 and pPL2 alleviate both of these concerns.

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It is evident from the above results and discussion that subject invention provides a number of advantages. The construction and characterization of the first single step site specific integration vectors for use in *L. monocytogenes* as described herein furthers the genetic tools available for the study of this pathogen. For example, the subject vectors and methods allow more facile strain construction than historic methods and are widely useful in various strains used to study the intracellular life cycle of *L. monocytogenes*. Furthermore, the subject invention provides important new tools for the production of vaccine preparations.

One limitation with plasmid-based expression of recombinant proteins in *L. monocytogenes* is the stability of the plasmids *in vivo* (*i.e.* in the host animal) without selection. Additionally, chromosomal construction of strains expressing foreign antigens is time consuming. The subject vectors and methods of use alleviate both of these concerns. As such, the present invention represents a significant contribution to the art.

All publications and patent application cited in this specification are herein incorporated by reference as if each individual publication or patent application were specifically and individually indicated to be incorporated by reference. The citation of any publication is for its disclosure prior to the filing date and should not be construed as an admission that the present invention is not entitled to antedate such publication by virtue of prior invention.

Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it is readily apparent to those of ordinary skill in the art in light of the teachings of this invention that certain changes and modifications may be made thereto without departing from the spirit or scope of the appended claims.

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